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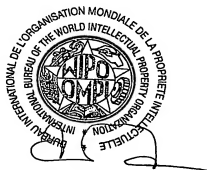
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Applicant's or agent's file reference (if desired) (12 characters maximum) **RS101** Box No. I TITLE OF INVENTION MEMBER FOR VASCULAR SEALING Box No. II APPLICANT This person is also inventor Name and address: (Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country. The country of the address indicated in this Box is the applicant's State (that is, country) of residence if no State of residence is indicated below.) Telephone No. Facsimile No. SCHNYDER, Guido Chemin de Champ Pallet 4 Teleprinter No. CH - 1801 Le Mont-Pélerin Applicant's registration No. with the Office State (that is, country) of nationality: State (that is, country) of residence: CH CH This person is applicant for the purposes of: all designated States all designated States except the United States of America the United States of America only the States indicated in the Supplemental Box FURTHER APPLICANT(S) AND/OR (FURTHER) INVENTOR(S) Name and address: (Family name followed by given name; for a legal entity, full official designation. The address must include postal code and name of country. The country of the address indicated in this Box is the applicant's State (that is, country) of residence if no State of residence is indicated below.) This person is: applicant only ROUVINEZ, Gilles applicant and inventor ch. de la Crétaz 25 inventor only (If this check-box is marked, do not fill in below.) CH - 1822 Chernex Applicant's registration No. with the Office State (that is, country) of nationality: State (that is, country) of residence: CH CH This person is applicant all designated all designated States except the United States of America the United States of America only the States indicated in the Supplemental Box for the purposes of: Further applicants and/or (further) inventors are indicated on a continuation sheet. Box No. IV AGENT OR COMMON REPRESENTATIVE; OR ADDRESS FOR CORRESPONDENCE The person identified below is hereby/has been appointed to act on behalf of the applicant(s) before the competent International Authorities as: common representative agent Name and address: (Family name followed by given name; for a legal entity, full official designation.

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Sheet No. ...3...

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| sequence listings and/or tables related thereto) : 13 | 3. ☐ original general power of attorney : 4. ☐ copy of general power of attorney; reference number, | | | | |
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MEMBER FOR VASCULAR SEALING

Field of the invention

The present invention relates to closing apertures in body tissues. More specifically, the present invention relates to sealing members such as a staple, clip, snap or rivet for scarring such apertures.

Background

. It frequently happens that portions of internal body tissue need to be sealed together. Often this need is a 10 result of а cardiac or peripheral vascular catheterization. The art of sealing body tissues will therefore be discussed with a particular emphasis on apertures resulting of such interventions. 15 Cardiac or peripheral vascular catheterizations are well known procedures that typically involve the making of a puncture in the femoral, radial or brachial artery to allow catheter insertions for diagnosis or treatment of cardiovascular or peripheral vascular diseases. After diagnostic and/or interventional catheterizations, the 20 puncture formed by the insertion of the catheter must be closed following removal of the catheter. The puncture opening in the artery has a typical size in the range of 4-6 French for diagnostic procedures and in the range of 25 6-15 French for interventional procedures. Traditionally, manual or mechanical compression are applied to puncture sites for at least 20 minutes and up to 6 hours after removal of the catheter. Other traditional methods for sealing the puncture site include the use of thrombotic or collagen plugs, patches, or other suturing methods. 30

In particular, patients who have had a femoral artery puncture should remain at strict bed rest, sometimes with a heavy sandbag on their groin for several hours, to ensure adequate hemostasis.

Traditional methods of hemostasis, as described above, 5 following a femoral artery access have many pitfalls. Patients have to remain on their back for many hours having their leg with the access site stretched, which is felt by many patients as a great discomfort, often 10 greater than the entire interventional procedure. Furthermore, the weight of a sandbag on the femoral artery often causes the lower leg to tingle or go numb. In addition, the longer it takes to obtain secure sealing of the wound (up to 24 hours), the higher the risk of 15 local complications such as hematoma, false aneurysms, local orsystemic infections and/or acute occlusions. This makes wound site management the longer critical care item involving additional costs, greater patient discomfort, and increased risk of complications.

20 Surgical stapling instruments have been proposed to solve some of the aforementioned problems associated with vascular procedures. U.S. Pat. No. 5,709,335 discloses a wholly distal surgical stapler for attaching a tubular vessel having two untethered ends. This stapler 25 is especially useful for making the primary permanent anastomotic connection of a bypass vein to a coronary artery or to the aorta. This stapler needs to temporarily placed within the tubular vessel (e.g., vein or artery). Such staplers are useful for stapling a graft vein or the like. However, they are inappropriate when 30 the entirety of the tubular tissue is not accessible,

such as during vascular sealing following cardiac or peripheral vascular catheterizations.

Another example of a surgical stapling instruments is found in U.S. Pat. No. 5,695,504 (Gifford et al.), which discloses a stapler to perform end-to-side anastomosis 5 between a graft vessel and the wall of a target vessel. The end of a graft vessel has to be passed through an inner sleeve of the stapler until the end of the vessel extends from the distal end of the stapler. The distal end of the graft is then permanently stapled to the wall 10 of the target vessel. Such staplers are useful attaching two tubular tissues together. However, they are inadequate for sealing vascular punctures, such as those created to perform cardiac or peripheral 15. catheterizations.

In general, staples used during cardiovascular surgery, and in particular staples of the type disclosed in U.S. Pat. No. 5,709,335 and 5,695,504, are known to be made out of durable materials such as for example titanium or stainless steel, which will outlive patients.

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Pat. No. 6,506,210 (assigned to AngioLink Corporation) discloses a wound site management and wound closure system involving a slightly different stapler. The staple does not require intraluminal delivery and is appropriate for sealing vascular punctures, such as those to perform cardiac or peripheral catheterizations. The assignee's system (EVS $^{\infty}$) involves a staple made of titanium, a biocompatible material with appropriate mechanical properties to allow efficient sealing of the puncture site. The assignee has emphasized the importance of a permanent durable material, such as

titanium, permitting X-ray puncture localization for subsequent interventions (Transcatheter Cardiovascular Therapeutics, Washington: September, 2002).

The use of inert durable materials such as titanium, stainless steal or ceramic, which do not dissolve during the lifetime of the patient, results in the implant of a permanent foreign body. This permanent implant may create a chronic irritation of the tissue surrounding the staple.

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10 Another drawback is the potential damaging effect of an external compression that can trap the underlying vessel between such a rigid permanent staple and the head of the femur. A laceration of the underlying vessel could therefore be caused by shocks, for example during falls 15 which are common in the case of the many debilitated patients undergoing cardiac or peripheral catheterizations.

Puncture of the femoral artery is ideally performed at the level of the common femoral artery, because of its 20 relatively large size and compressability. The latter depends on its course over bone, against which it can be readily compressed to achieve hemostasis. If the puncture is too proximal, the external iliac artery may be entered, increasing the risk of retroperitoneal 25 hemorrhage; if the puncture is too distal, either the profunda femoral artery or the superficial femoral artery can be punctured, with a risk of local complications such as vessel laceration, pseudoaneurysm, arteriovenous fistula, thrombosis, or excessive bleeding.

The anatomy and in particular the length of the common femoral artery have been quantified to allow optimal puncture thereof. It has been found that the ideal puncture site is located in the area overlapping the 5 inner quadrant of the femoral head, accurately predicts access in the common femoral artery whose length ranges from 0 to 11cm (mean: 6.7cm). This limits the remaining accessible segment of the common femoral artery to a length of about 2.0cm (Schnyder, G. 10 al., in Catheterization and Cardiovascular Interventions, vol. 53, pp. 289-295 [2001]). Since the staple described in U.S. Pat. No. 6,506,210 diameter of 3-4mm when fully expanded during implantation, it follows that such a staple can only be used a limited number of times (two to three) at the same 15 location. This is problematic for treating patients with extensive vascular disease, who require a plurality of interventions.

Summary of the invention

20 It is an object of the present invention to reduce the of injuries orother complications following application of the above described prior art sealing members. The present invention relates to a member for urging together two or more portions of body tissue and maintaining these portions together until 25 they secured together by scarring thereof. According to the invention, the member is made of a material selected from at least one of metals, alloys and ceramic compounds thereof such as oxides, which material is bioresorbable 30 and/or biodegradable.

The above portions of body tissue may form a wound, such as a puncture resulting from a catheter-based intervention. In the content of the present invention, any puncture is contemplated, accidental or intentional.

- 5 The sealing member of the present invention can be used in and around the femoral, radial, and brachial arteries after coronary, cardiac or peripheral vascular procedures. The sealing member can also be used for vessel or any tube like body-part clamping or occlusion, 10 soft-tissue anchoring, tendon and artery joining, meniscal repair, thoracic lung closure, heart repair, endoscopic procedures, esophageal repair, laparoscopy, skin or epidermal wound closure and general tissue closure.
- A bioresorbable material is a material that is transformed, when present in a body tissue, into smaller elements such as colloidal particles with the newly formed elements remaining in the body as traceable elements containing for example titanium, zirconium, niobium oxide, tantalum, silicon and lithium or compounds thereof.

A biodegradable material is a material that is transformed, when present in body tissue, into smaller elements - such as soluble salts - with the newly formed elements either remaining in the surrounding tissue as fine undetectable precipitates or dissolving and being ultimately eliminated from the body. These elements include for example magnesium, zinc, sodium, potassium, calcium, iron and manganese salts or compounds thereof.

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The nature of the materials used in the members of the present invention - bioresorbable / biodegradable - are opposed to the prior art biocompatible or bioabsorbable members, which both permanently remain in the local body tissue without undergoing any major structural changes. The biocompatible materials remain inert and do not trigger any tissue counter-reaction, while bioabsorbable materials ultimately are incorporated permanently into the surrounding tissue.

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The sealing member of the present invention may be used to close an artery or vein following a diagnostic or interventional procedure. More generally, the member may be used for any tissue repair.

As opposed to prior art permanent staples, the sealing
15 member according to the inventions is present in the
human body for a limited period of time sufficient to
secure scarring, thereby tremendously reducing the risk
of subsequent vessel injuries by external compression and
permitting unlimited repetitive use for future
20 interventions.

implantable, bioresorbable and/or biodegradable sealing member may comprise a combination of materials which dissolve in the human body without any harmful effects on the person that wears the member. materials of a sealing member can be a combination of metals, polymers or mixture thereof or any substances such that degradation products originating from the sealing member in the form of particles of at least one of soluble salts, fine particles (e.g. $0.1 \mu m$ - $50\mu m$) and/or colloidal particles (e.g. $5nm-0.1\mu m$).

The sealing member is advantageously made of materials which can undergo adequate plastic deformation (to enable insertion of the member with negligible elastic recoil) and strong mechanical properties so as to secure the wound site despite shear forces generated by blood flow within the vessel and by the surrounding tissue when the patient is moving (i.e. walking, climbing stairs, etc...).

It will be appreciated throughout the following description that the members of the present invention are made of any bioresorbable and/or biodegradable material that fulfills the above required properties of deformability and mechanical resistance.

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Such materials have previously been used to manufacture vessel wall supports or stents, as described in U.S. Pat. 15 No. 6,287,332 (assigned to Biotronik Mess-Therapiegeraete GmbH & Co.), the disclosure of which is hereby incorporated by way of reference. Such stents are used to minimize inflammatory reaction so as to reduce production of scar material upon implantation, 20 whereby instent restenosis or renarrowing of previously treated vessel segment by scar material is prevented. Surprisingly, sealing members made of these bioresorbable and/or biodegradable materials according to the invention do not prevent secure scarring but permit 25 adequate healing.

The material of the member can be of a combinations of metals which can dissolve in the body without significantly forming bio-incompatible decomposition products. The material may dissolve at a rate in the range from 0.1 to 5 mg/day, in particular from 0.5 to 2mg/day. A sealing member made of this material may be

entirely dissolved within 1 to 300 days, in particular 5 to 100 days, such as 10 to 50 days.

Such a temporary sealing member combines the mechanical properties of metals with the bioresorbability polymer-based materials. Immediate mechanical sealing of an access site can be achieved with this non-permanent The sealing member may be entirely sealing member. dissolved after complete scarring of the access site or after sufficient scar material has been formed to permit complete sealing even upon dissolution or resorption of the sealing member. The sealing member should retain it's mechanical properties to maintain the body tissues urged together for at least 1 day, in particular for at least 3 days.

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In a first embodiment of the invention, the sealing 15 is made of a metal alloy suitable for member biocompatible decomposition, as explained in below. Consequently, in this embodiment, the metal alloy consists essentially of a combination of materials that 20 will decompose in the body comparatively rapidly - within a period of days or weeks or months, but preferably no more than 12 months - forming harmless products.

To obtain a substantially uniform decomposition, such an alloy may comprise a component A which covers itself with a protective oxide coat. This component A can be selected from one or several metals of consisting of magnesium, titanium, zirconium, niobium, silicon. Moreover, tantalum, zinc to obtain or substantially uniform dissolution of this oxide coat, a component B, that possesses sufficient solubility in interstitial fluids or blood - such as lithium, sodium,

potassium, calcium, iron or manganese - can be added to the alloy.

Suitable metals for the alloy include metals that are naturally present in the human body (magnesium, zinc, sodium, potassium, calcium, iron and manganese) or that are nontoxic (titanium, zirconium, niobium, tantalum, silicon and lithium). The combination of a passivating and a soluble component ensures a timely and uniform decomposition into biocompatible breakdown products. The decomposition rate can be set by the ratio of the two components.

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The alloy can be such that the decomposition products are soluble salts, in particular sodium, potassium, calcium, iron or zinc salts, or non-soluble decomposition products, such as titanium, tantalum or niobium oxide, in the form of colloidal particles. The decomposition rate is advantageously adjusted by the composition so that gases, when formed, dissolve physically without forming any macroscopic gas bubbles. For example, hydrogen gas evolves during the decomposition of lithium, sodium, potassium, magnesium, calcium or zinc salts.

For instance, an alloy of lithium and magnesium, can be used as a possible alloy which is however optimized with a view to increase fatigue durability for the field of application mentioned above. The weight ratio magnesium/lithium can be of the order of 60/40, fatigue durability being increased by the addition of further components, such as zinc, or by gassing by hydrogen. Also, special melting and forging methods can be used to increase the fatigue durability.

A sodium-magnesium alloy may be used to make the sealing member. Since sodium hydroxide as a decomposition product possesses a high solubility, this alloy can brake down without voluminous encrusting. Sodium dissolves and magnesium hydroxide forms a fine precipitate which may deposit without risk in the surrounding tissue.

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Another decomposable combination of metal materials is a zinc-titanium alloy, with a percentage by weight of titanium in the range of 0.1% to 1%. This combination precludes the comparatively strong crystalline growth of zinc during use, which would cause a comparatively brittle and fragile behavior of the sealing member. The addition of titanium leads to the formation of a $Zn_{15}Ti$ phase at the crystal boundaries, which precludes any further crystalline growth. This reduction of the grain size generally improves the ductility, in particular it increases the elongation at rupture.

If gold is added to this alloy at a percentage by weight of 0.1% to 2%, a further reduction of the grain size is achieved when the material cures. This further improves the tensile strength of the material.

In another embodiment of the invention, the sealing member comprises a support body and a local electrode for use as an electrochemical device. The support body can be made of a substantially pure metal. Usually, the local electrode is made of a second metal and is in contact with the support body. This local electrode can be a coat on the sealing member or is fixed onto the sealing support body by electroplating or by laser welding. The contact between the support body and the local electrode produces a contact voltage and a resulting current that

leads to active degradation of the sealing member. The degradation rate and thus the decomposition rate of the sealing member can be controlled by the size of the local electrode and by the selection of the metals of the sealing member.

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Detailed description

The invention will be further explained in the following Exemples:

Example 1

10 A bioresorbable and/or biodegradable sealing member according to the invention can be made from an allow containing zinc as the component A and calcium as the component B. The weight ratio that zinc bears to calcium amounts to 25/1. This Zn-Ca alloy forms soluble salts as 15 degradation products, such as calcium hydroxide which possesses such a high solubility that the solubility product is not transgressed during slow decomposition several weeks ormonths. This hydroxide transported in dissolved form by interstitial fluids or 20 blood and is metabolized.

To improve the mechanical properties of the sealing member, such as ductility, hardness and tensile strength, suitable alloy constituents can be added in low concentrations. For instance, phosphorus may be added to the alloy in an amount of the order of a few percents.

Example 2

A bioresorbable and/or biodegradable metal sealing member acting as a local electrochemical device according to the invention can comprise a support body and a local

electrode. The support body is made of substantially pure zinc which dissolves - as electroplating tests show - without production of gases and without the formation of oxide at currents of some milliamps. The local electrode is made of gold and is in contact with the zinc support body. This local gold electrode is fixed onto the sealing support body by electroplating or by laser welding. The contact between the zinc support body and the local gold electrode produces a contact voltage and a resulting current that leads to active degradation of the sealing member. The exchange current as a whole is determined by the size of the gold electrode. The degradation rate and thus the decomposition rate of the sealing member can be adjusted by the size of the local gold electrode.

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Tests have shown that an exchange current arises between the sealing member's support body and the local electrode after few minutes and remains constant for several days in such a sealing member. Hence, a constant decomposition rate can be attained and a member of 10 mg will dissolve within approximately 30 to 40 days at a corrosion current of 10 μA.

Example 3

A bioresorbable metal sealing member according to the invention can be made form a ZnTi alloy with a Ti weight percentage of 0.1% to 1%. In a further embodiment of this example, a precious metal in the form of gold can be added at a weight percentage of 0.1% to 2%, the Ti weight percentage remaining unchanged so that the member consists of a ZnAuTi alloy. These two alloys also exhibit a biocompatible decomposition behavior and are thus regarded as bioresorbable sealing members.

CLAIMS

- 1. A member for urging together two or more portions of body tissue and maintaining said portions together until said portions are secured together by scarring thereof, wherein said member is made of a material selected from at least one of metals, alloys and ceramic compounds thereof, such as oxides, which material is bioresorbable and/or biodegradable.
- 2. The member of claim 1, which is a staple, clip, snap 10 or rivet.
 - 3. The member of claim 1 or 2, wherein said material is a metal alloy containing: a first component which covers itself with a protective oxide coat; and a second component which ensure sufficient dissolution of the oxide coat.

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- 4. The member of claim 3, wherein the first component comprises at least one metal selected from magnesium, titanium, zirconium, niobium, tantalum, zirc and silicon and the second component comprises at least one metal selected from lithium, sodium, potassium, manganese calcium and iron.
- 5. The member of claim 3 or 4, wherein the components of the metal alloy are selected such that corrosion products originate therefrom in the form of soluble salts, fine particles or colloidal particles or a mixture thereof.
- 6. The member of claim 3, 4 or 5, wherein the alloy contains zinc as a corrosion-inhibiting component.
- 7. The member of claim 6, wherein the alloy contains zinc and calcium.

- 8. The member of claim 7, wherein the alloy has a zinc/calcium weight ratio of at least 21/1.
- 9. The member of claim 3, 4, or 5, wherein the alloy contains sodium and magnesium.
- 5 10. The member of claim 1 or 2, wherein the bioresorbable and/or biodegradable material is an alloy of zinc and titanium.
 - 11. The member of claim 10, wherein the zinc-titanium alloy has a weight percentage of titanium of 0.1% to 1%.
- 10 12. The member of claim 11, wherein an amount of 0.1 to 2 weight% gold is added as a further component to the zinc titanium alloy.
 - 13. The member of claim 1 or 2, wherein the bioresorbable and/or biodegradable sealing member comprises a support
- 15 body made of a substantially pure first metal and a local electrode made of a second metal which is in contact with the support body to produce a contact voltage and a resulting current that leads to active degradation of the sealing member.
- 20 14. The member of claim 13, wherein the local electrode is a coat on the support body.
 - 15. The member of claim 13, wherein the local electrode is a metal part attached to the support body.
- 16. The member of claim 13, 14 or 15 wherein the support body consists essentially of zinc.
 - 17. The member of claim 13, 14 or 15 wherein the local electrode consists essentially of a precious metal.

- 18. The member of claim 14, wherein said coat is deposited by electroplating or sputtering.
- 19. The member of any preceding claim, wherein the sealing member is made of a phosphorus-containing alloy.
- 5 20. The member of any preceding claim, which is a hydrogen-treated alloy.
- 21. The member of any preceding claim, which is made of an alloy which during use corrodes at such a rate that gases arising during corrosion physically dissolves in a body fluid to which the alloy is exposed.

ABSTRACT

An implantable, bioresorbable and/or biodegradable sealing member, such as a staple, clip, snap or rivet, is used for clamping vessels, vessel side-branches, aneurysms, or any other tube like body-parts or for sealing vascular access sites and wound site management. The implantable, bioresorbable and/or biodegradable member comprises a combination of materials which dissolve or degrade in the human body without any harmful effects on the person that wears the member.

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